
 (Label* NEAR packag*) NEAR device* NEAR steril*



SmartSearch Filters

FDA Regulation (21 CFR/Guidance) 

Select value



FDA Agency (CBER/CDRH/CDER)

Devices (23)



FDA Search Results Shown Below 



[ FDA Agency (CBER/CDRH/CDER) ="Devices"]

23 results for '(Label* NEAR packag*) NEAR device* NEAR steril* '

Click to Sort SmartSearch Results



[Subpart E - Other Exemptions](#)

**Medical devices; processing, labeling, or repacking.
Sec. 801.150(e)**



[Subpart E - Cardiovascular Surgical Devices](#)

**Apical closure device.
Sec. 870.4510(b)(3)**



[Subpart B - Diagnostic Devices](#)

Endoscopic transhepatic venous access needle.
Sec. 876.1050(b)(4)



[Subpart E - Surgical Devices](#)

Endoscopic electro surgical clip cutting system.
Sec. 876.4310(b)(5)



[Subpart F - Therapeutic Devices](#)

Metallic biliary stent system for benign strictures.
Sec. 876.5011(b)(5)



[Subpart F - Therapeutic Devices](#)

Nonimplanted nerve stimulator for functional abdominal pain relief.
Sec. 876.5340(b)(6)

1 2 3 4 »



storage. is there anything else you would like me to clarify



11:45

Which label requirement can be applied at the package level only not individual tray

11:46

The expiration date is the only label requirement that can be applied at the package level. All other label requirements, such as the unique identifier and lot number, must be applied to each individual tray within the package.

11:46

What id there is not enough space on the tray

11:51

If there is not enough space on the tray for all the required label information, then alternative methods such as side panel labeling or multi-layer labeling may be used. However, it is important to ensure that all necessary information is still easily visible and accessible for proper tracking and identification purposes.

✎ Reply to GPT-3

**Email**